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NEWS				CA/CAplus enhanced with printed CA page images from
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NEWS	13	SEP	17	CAplus coverage extended to include traditional medicine
				patents
NEWS	14	SEP	24	EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS	15	OCT	02	CA/CAplus enhanced with pre-1907 records from Chemisches
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NEWS	16	OCT	19	BEILSTEIN updated with new compounds
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NEWS		NOV		ICSD reloaded with enhancements
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NEWS		DEC		USPATOLD added to additional database clusters
NEWS	23	DEC	17	IMSDRUGCONF removed from database clusters and STN
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NEWS	28	DEC	17	STN Viewer enhanced with full-text patent content
				from USPATOLD
NEWS		JAN		STN pricing information for 2008 now available
NEWS	30	JAN	16	CAS patent coverage enhanced to include exemplified
				prophetic substances
NEWS	31	JAN	28	USPATFULL, USPAT2, and USPATOLD enhanced with new
MIDITO	2.0	7711	0.0	custom IPC display formats
NEWS		JAN		MARPAT searching enhanced
NEWS	33	JAN	28	USGENE now provides USPTO sequence data within 3 days
NUMBER	2.4	JAN	20	of publication
NEWS	34	UAN	28	TOXCENTER enhanced with reloaded MEDLINE segment

NEWS 35 $\,$ JAN 28 $\,$ MEDLINE and LMEDLINE reloaded with enhancements NEWS 36 $\,$ FEB 08 $\,$ STN Express, Version 8.3, now available

NEWS EXPRESS FEBRUARY 08 CURRENT WINDOWS VERSION IS V8.3, AND CURRENT DISCOVER FILE IS DATED 24 JANUARY 2008

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- => S Propyene glycol (S) Tonicity (S) (liquid formulation) AND pd<=20030708 2 FILES SEARCHED...
- L1 0 PROPYENE GLYCOL (S) TONICITY (S) (LIQIUD FORMULATION) AND PD<=20 030708
- => S (Propylene glycol) (S) Tonicity (S) (liquid formulation) AND pd<=20030708 2 FILES SEARCHED...
- .2 0 (PROPYLENE GLYCOL) (S) TONICITY (S) (LIQUID FORMULATION) AND PD<=20030708</p>
- => S (Propylene glycol) (P) Tonicity (P) (liquid formulation) AND pd<=20030708 2 FILES SEARCHED...
- L3 0 (PROPYLENE GLYCOL) (P) TONICITY (P) (LIQUID FORMULATION) AND PD<=20030708
- => S (Propylene glycol) (S) (liquid formulation) AND pd<=20030708 2 FILES SEARCHED...
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PROCESSING COMPLETED FOR L4

17 DUP REM L4 (4 DUPLICATES REMOVED) ANSWERS '1-3' FROM FILE BIOSIS ANSWERS '4-17' FROM FILE CAPLUS

=> D ti 15 1-17

- L5 ANSWER 1 OF 17 BIOSIS COPYRIGHT (c) 2008 The Thomson Corporation on STN DUPLICATE 1
- TI A STABILITY STUDY OF CLINDAMYCIN HYDRO CHLORIDE AND CLINDAMYCIN PHOSPHATE SALTS IN TOPICAL FORMULATIONS.
- L5 ANSWER 2 OF 17 BIOSIS COPYRIGHT (c) 2008 The Thomson Corporation on STN DUPLICATE 2
- TI ASPIRIN DEGRADATION IN MIXED POLAR SOLVENTS.
- L5 ANSWER 3 OF 17 BIOSIS COPYRIGHT (c) 2008 The Thomson Corporation on STN
- TI Diffusion of herbicides through plastic film.
- L5 ANSWER 4 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Stabilizing biomolecules in liquid formulations
- L5 ANSWER 5 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Stable viscous liquid formulations of amlexanox for the prevention and treatment of mucosal diseases and disorders
- L5 ANSWER 6 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Transdermal Delivery of Highly Lipophilic Drugs: In Vitro Fluxes of Antiestrogens, Permeation Enhancers, and Solvents from Liquid Formulations
- L5 ANSWER 7 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Pharmaceutical compositions containing chelates and reducing agents with improved stability
- L5 ANSWER 8 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Stable liquid formulations of high vitamin E content
- L5 ANSWER 9 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Antiperspirant formulation for porous applicator
- L5 ANSWER 10 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Nebulizer-compatible liquid formulations for pulmonary delivery of glucocorticoids: pre-formulation studies
- L5 ANSWER 11 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Pharmaceutical compositions containing lamivudine and a preservative
- L5 ANSWER 12 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Stable particle in liquid formulations comprising sugar glass
- L5 ANSWER 13 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Compositions comprising an HIV protease inhibitor such as VX 478 and a water soluble vitamin e compound such as vitamin E-TPGS
- L5 ANSWER 14 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Tastemasked liquid pharmaceuticals containing sugars and hydrogenated maltose and polyhydroxy alcohols
- L5 ANSWER 15 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TI Stabilized isothiazolone liquid formulation

- L5 ANSWER 16 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- TΤ Stabilized aqueous liquid formulations of phytase and their use in feed preparation for monogastric animals
- ANSWER 17 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
- Metal-acid complexes with members of the tetracycline family. II. TT Development of stable preconstituted parenteral formulations

=> D ibib abs 15 1-17

ANSWER 1 OF 17 BIOSIS COPYRIGHT (c) 2008 The Thomson Corporation on STN DUPLICATE 1

ACCESSION NUMBER: 1984:337147 BIOSIS

DOCUMENT NUMBER: PREV198478073627; BA78:73627

TITLE: A STABILITY STUDY OF CLINDAMYCIN HYDRO CHLORIDE AND CLINDAMYCIN PHOSPHATE SALTS IN TOPICAL FORMULATIONS. AUTHOR(S): MIGTON J M [Reprint author]; KENNON L; SIDEMAN M;

PLAKOGIANNIS F M

CORPORATE SOURCE: DIV OF PHARMACEUTICS AND INDUSTRIAL SCI, ARNOLD AND MARIE

SCHWARTZ COLL OF PHARMACY AND HEALTH SCI, LIU, 75 DEKALB

AVE, BROOKLYN, NY 11201, USA SOURCE: Drug Development and Industrial Pharmacy, (1984)

Vol. 10, No. 4, pp. 563-574.

CODEN: DDIPD8, ISSN: 0363-9045.

DOCUMENT TYPE: Article FILE SEGMENT: LANGUAGE: ENGLISH

The stability of clindamycin hydrochloride and clindamycin phosphate [used in the treatment of acne vulgaris] was studied in topical liquid

formulations prepared with the following solvents: solvent a (70%

isopropanol, 10% propylene glycol and 20% water);

solvent B (48% isopropanol, polyoxyethelene ethers, acetone, salicylic acid and allantoin); solvent C (40% alcohol, acetone, polysorbate 20,

fragrance and water); and standard (50% isopropyl alcohol,

propylene glycol and water) in glass and plastic containers at 25°, 40°, and 50° C. In general,

better stability was obtained in glass containers than in plastic

containers. At 25° C both the clindamycin hydrochloride and

phosphate formulations in solvent B showed poorer stability than in the other solvents irrespective of the type of container, while formulations in solvent C showed the best stability. The effect of the pH on the stability of the formulations was determined. At pH values below 4 the stability of all formulations decreased.

ANSWER 2 OF 17 BIOSIS COPYRIGHT (c) 2008 The Thomson Corporation on STN DUPLICATE 2

ACCESSION NUMBER: 1984:309646 BIOSIS

DOCUMENT NUMBER: PREV198478046126; BA78:46126

ASPIRIN DEGRADATION IN MIXED POLAR SOLVENTS. TITLE: AUTHOR(S): CHANG R-K [Reprint author]; WHITWORTH C W CORPORATE SOURCE: COLL PHARM, UNIV GA, ATHENS, GA 30602, USA SOURCE:

Drug Development and Industrial Pharmacv, (1984)

Vol. 10, No. 3, pp. 515-526.

CODEN: DDIPD8. ISSN: 0363-9045.

DOCUMENT TYPE: Article FILE SEGMENT: BA

LANGUAGE: ENGLISH

Degradation studies were conducted 0.2% w/v [wt/vol] aspirin [an AB antipyretic, antiinflammatory and analgesic agent] liquid formulation in a wide range of water-propylene

glycol mixtures and water-triethylene glycol diacetate mixtures at temperatures. The effect of a surfactant, polyoxyethylene (20) sorbitan monolaurate, on aspirin stability was also investigated. There was a linear relationship between water content and degradation rate constants. The surfactant increased aspirin degradation in all formulations. Formulations containing the higher concentration of the surfactant showed the greater aspirin decomposition.

L5 ANSWER 3 OF 17 BIOSIS COPYRIGHT (c) 2008 The Thomson Corporation on STN ACCESSION NUMBER: 1964:32033 BIOSIS

DOCUMENT NUMBER: PREV19644500032037; BA45:32037

TITLE: Diffusion of herbicides through plastic film.

AUTHOR(S): BRIDGES, W. R.; SANDERS, HERMAN O.

CORPORATE SOURCE: U. S. Bur. Sport Fish and Wildlife, Denver, Colo., USA

SOURCE: PROGR FISH CULT, (1963) Vol. 25, No. 4, pp. 213-214.

DOCUMENT TYPE: Article FILE SEGMENT: BA

LANGUAGE: Unavailable

ENTRY DATE: Entered STN: May 2007

Last Updated on STN: May 2007

AB Laboratory tests with various herbicides and polyethylene and saran film demonstrated that herbicides will diffuse through these materials in aquatic situations. Tests with a liquid formulation of the propylene glycol butyl ether esters of 2,4-D and polyethylene bags of 0.003-in. thickness, revealed that when 10 mg. of the herbicide was added to 5 1. of water in the bag and the bag was immersed in 10 1. of water, and equilibrium of herbicide in the water inside and outside the bag was reached after 96 hours. Similarly conducted tests with polyethyl ether in an indicated that it is an

onducted tests with polyvinyl chloride film indicated that it is an effective barrier. Diffusion through the vinyl film did not occur in tests using the propylene glycol butyl ether esters of 2,4-D and the butoxyethanol ester of silvex. ABSTRACT AUTHORS: W. R. Bridges

L5 ANSWER 4 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:428730 CAPLUS

DOCUMENT NUMBER: 137:10994

TITLE: Stabilizing biomolecules in liquid formulations
INVENTOR(S): Cowan, Siu Man L.; McGinnis, Vincent; Palmer, Donna

T.; Risser, Steven M.; Brody, Richard S.

PATENT ASSIGNEE(S): Battelle Memorial Institute, USA

SOURCE: PCT Int. Appl., 20 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

				KIND DATE				APPLICATION NO.						DATE			
WO 200				A2 20020606 A3 20021031				WO 2	001-	US48		20011030 <					
WO 200	2043/	50		A3		2002	1031										
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	co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FΙ,	GB,	GD,	GE,	GH,	
	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	
	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,	
	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	ΤJ,	TM,	TR,	TT,	TZ,	UA,	
	UG,	UZ,	VN,	YU,	ZA,	ZW											
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	IE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	BJ,	CF,	CG,	CI,	CM,	GA,	GN,	
	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG									

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A1 20020606 CA 2001-2430137
A 20020611 AU 2002-36641
A2 20030917 EP 2001-986180
                                                                    20011030 <--
     CA 2430137
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     AU 2002036641
     EP 1343521
                                                                     20011030
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     JP 2004526674
                          T
                               20040902 JP 2002-545720
                                                                      20011030
                                20020801 US 2001-20798
     US 2002102218
                          A1
                                                                      20011130 <--
     US 2002110524
                         A1 20020815 US 2001-20799
B2 20061128
                                                                      20011130 <--
    US 7141542
                        A 20050930 NZ 2001-526135
A 20040504 MX 2003-PA4883
A1 20050526 US 2004-15201
     NZ 526135
                                                                      20011130
     MX 2003PA04883
US 2005112092
                                                                      20030530
                                                                      20041217
PRIORITY APPLN. INFO.:
                                              US 2000-250491P
                                                                  P 20001201
                                                                  W 20011030
                                              WO 2001-US48834
                                              US 2001-20798
                                                                  B1 20011130
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AB The invention is directed to a stable formulation of a biol. active protein useful for aerosol delivery to the respiratory tract of a patient in need of treatment comprising; (a) a carrier liquid comprising from about 10 % to from about 100 % V/V water and from about 0 % to from about 90 % V/V of an organic liquid; (b) a biol. effective amount of a protein suspended

or

dissolved in a carrier liquid; and (c) a stabilizing effective amount of a derivatized carbohydrate stabilizing agent suspended or dissolved in said carrier liquid The stable formulations of the invention may optionally contain about 0.1 % to about 5.0 % weight/volume of a pharmaceutically acceptable excipient. In an ethanol-water (80:20) carrier liquid the preferred stabilizer for insulin is C12-glucose, while in a totally aqueous carrier liquid the preferred stabilizer is C8 glucose or C8 trehalose.

L5 ANSWER 5 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2002:575762 CAPLUS

ACCESSION NUMBER: 2002:5/5/62 C

DOCUMENT NUMBER: 137:129916

TITLE: Stable viscous liquid formulations of amlexanox for the prevention and treatment of mucosal diseases and

disorders

INVENTOR(S): Jacob, Jeremy

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 22 pp. CODEN: USXXCO

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE .	APPLICATION NO.	DATE
US 2002103219	A1	20020801	US 2001-971562	20011004 <
PRIORITY APPLN. INFO.:			US 2000-238175P P	20001005
3 D D 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			3.1 1.1 6 3 1.1	

AB Present invention concerns stable viscous liquid formulations of amlexanox for the prevention and treatment of mucosal diseases and disorders. The mucoadhesive of the present invention may be a linear or crosslinked polymer such as polyacrylic acid, hydroxyalkyl cellulose, dextran sulfate, and etc. An object of the present invention is to provide a convenient and effective dosage form for Amlexanox in the treatment of skin mucous disorders. This form allows for an ED of the pharmaceutical to be applied to the lesions being treated over an extended period. Thus, a viscous, mucoadhesive aqueous composition contained water 91.26, KoH 0.60, benzyl alc.

1.50,

Polysorbate-60 0.05, Carbopol 971P 0.35, H3PO4 0.13, citric acid 0.05, saccharin sodium 0.40, amlexanox 0.50, and glycerin 5.20% by weight

L5 ANSWER 6 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

AUTHOR(S):

CORPORATE SOURCE:

SOURCE:

AB

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GT

2002:454133 CAPLUS

138:226493

Transdermal Delivery of Highly Lipophilic Drugs: In Vitro Fluxes of Antiestrogens, Permeation Enhancers, and Solvents from Liquid Formulations Funke, Adrian P.; Schiller, Roman; Motzkus, Hans W.; Guenther, Clemens; Mueller, Rainer H.; Lipp, Ralph Pharmaceutical Development, Schering AG, Berlin,

13342, Germany

Pharmaceutical Research (2002), 19(5),

661-668

CODEN: PHREEB: ISSN: 0724-8741 Kluwer Academic/Plenum Publishers

Journal

English

and AE 2 (II) (log P = 7.8) shall be delivered transdermally. Transdermal permeation of drugs, enhancers, and solvents from various fluid formulations were characterized by in-vitro permeation studies through excised skin of hairless mice. Furthermore, differential scanning calorimetry (DSC) measurements of skin lipid phase transition temps. were conducted. Transdermal flux of highly lipophilic drugs was extraordinarily enhanced by the unique permeation enhancer combination propylene glycol-lauric acid (9 & 1): steady-state fluxes of AE 1 and AE 2 were as high as 5.8 μg·cm-2·h-1 and 3.2 μg·cm-2·h-1, resp. This dual enhancer formulation also resulted in a marked increase in the transdermal fluxes of the enhancers. Furthermore, skin lipid phase transition temps. were significantly reduced by treatment with this formulation. Transdermal delivery of highly lipophilic drugs can be realized by using the permeation enhancer combination propylene glycol-lauric acid. The extraordinary permeation enhancement for highly lipophilic drugs by this formulation is due to mutual permeation enhancement of these two enhancers and their synergistic lipid-fluidizing activity in the stratum corneum.

Highly lipophilic basic drugs, the antiestrogens AE 1 (I) (log P = 5.82)

L5 ANSWER 7 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:247182 CAPLUS

DOCUMENT NUMBER: 134:271268

TITLE: Pharmaceutical compositions containing chelates and

reducing agents with improved stability

INVENTOR(S): Khanolkar, Jayant Eknath
PATENT ASSIGNEE(S): Procter & Gamble Co., USA

PATENT ASSIGNEE(S): Procter & Gamble Co., USA SOURCE: PCT Int. Appl., 36 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 4 PATENT INFORMATION:

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PAT	TENT :	NO.			KIN	D										DATE		
	2001	0229 AE, CR, HU, LU,	AG, CU, ID, LV,	AL, CZ, IL, MA,	A1 AM, DE, IN, MD,	AT, DK, IS, MG,	2001 AU, DM, JP, MK,	0405 AZ, DZ, KE, MN,	BA, EE, KG, MW,	WO 2 BB, ES, KP, MX,	000- BG, FI, KR, MZ,	JS26 BR, GB, KZ, NO,	402 BY, GD, LC, NZ,	BZ, GE, LK, PL,	CA GH LR PT	20000 , CH, , GM, , LS, , RO,	926 CN, HR, LT, RU,	
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HU	R: 2002	AT, IE, 0027	BE, SI, 97	CH,	DE, LV, A2	DK, FI,	ES, RO, 2003	FR, MK, 0128	GB, CY,	GR, AL	IT,	LI,	LU,	NL,	SE	, MC, 20000	PT,	
JP AU	2002 2003 7703 2002	5102 76	79		T B2			0318 0219		AU 2		7986	5			20000 20000 20020	926	
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AB The present invention pertains to improved stability of compns. that deliver drugs. These compns. have exceptional stability when used in various product forms including liquid elixirs placed into the mouth and eventually swallowed, or can be delivered via liquid-filled lozenges, metered liquid dosing devices, atomizers and liquid-releasing, edible capsules. Such compns. are particularly useful for treating symptoms associated with respiratory illnesses. Thus, a liquid formulation contained dextromethorphan 3.425, sodium

hexametaphosphate 0.050, propylene glycol 95.355,

sucralose 0.300, Pro-Sweet liquid-K 0.700, monosodium glycyrrhizinate 0.050, flavor 0.015, colorant 0.005, and sodium thiosulfate 0.100% by weight REFERENCE COUNT: 6 THERE ARE CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 8 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:895653 CAPLUS

DOCUMENT NUMBER: 136:25112

TITLE: Stable liquid formulations of high vitamin E content

INVENTOR(S): Crepeau, Michel Andre

PATENT ASSIGNEE(S): Aventis Animal Nutrition, S.A., Fr.

SOURCE: U.S., 3 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA		KIND DATE			APPLICATION NO.						DATE							
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US	6329	423			B1		2001	1211		US 2	2000-	5908	04		2	0000	609	<
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										US 2	-000	5908	04		A 2	0000	609	
										WO 2	2001-	EP39	52		A 2	0010	323	

A stable liquid vitamin E formulation having at least 60% vitamin E comprises water 0.5-3, potassium sorbate 0.05-0.15, propylene glycol 0.3-0.7, 1-propanol 15-20, polyethylene glycol 400 monooleate 12-17 and vitamin E oil 60-70% by weight The formulation is free of polyoxyethylene sorbitan monooleate and has a viscosity at 20° of <about 1000 cPs.

L5 ANSWER 9 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1999:748166 CAPLUS

DOCUMENT NUMBER: 131:341784

TITLE: Antiperspirant formulation for porous applicator INVENTOR(S):

Schamper, Thomas; Moghe, Bhalchandra; Barr, Morton L.; Wu, Ching-min Kimmy

PATENT ASSIGNEE(S): Colgate-Palmolive Company, USA

SOURCE: U.S., 11 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5989531	A	19991123	US 1998-191897	19981113 <
CA 2349167	A1	20000525	CA 1999-2349167	19991102 <
WO 2000028956	A1	20000525	WO 1999-US25570	19991102 <

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            CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,
            IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA,
            MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI,
            SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE,
            DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF,
            CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
    BR 9915298
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                                                                  19991102 <--
    EP 1128803
                         A1
                               20010905
                                          EP 1999-957489
                                                                  19991102 <--
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            IE, FI, RO
    HU 2001004518
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                                                                  19991102 <--
    HU 2001004518
                         A3
                               20030428
    AU 760372
                         B2
                               20030515
                                           AU 2000-15183
                                                                  19991102 <--
    ZA 2001003811
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                                           ZA 2001-3811
                                                                  20010510 <--
                         A
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                         Α
                                                                  20010514 <--
PRIORITY APPLN. INFO.:
                                           US 1998-191897
                                                               A 19981113
                                                               W 19991102
                                           WO 1999-US25570
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The invention comprises a liquid composition which provides a drier feel and AB reduced leakage when used with certain types of applicators, especially an applicator having a porous surface, which composition is made by combining an active phase and a silicone phase. The active phase is made by combining: (a) 10-70 % of a selected glycol; (b) 0.1-10 % of a nonionic emulsifier having an HLB greater than 8; (c) 0.01-30 % of a cosmetically active ingredient; and (d) 0-20 % of ethanol and/or isopropanol. The silicone phase is made by combining: (a) 0.1-10 % of a selected emulsifier; (b) 0-30 % of a non-volatile silicone; (c) 0-30 % of a volatile silicone; and (d) 0-25% of an organic emollient; provided that, (a) the silicone phase contains ≥ 10 % silicone; (b) the ratio of silicone phase to the active phase is in the range of 1:1 to 1:4; and (c) the composition is processed to maintain a viscosity in the range of 2,000-200,000 cP. A clear antiperspirant composition was made by combining dimethicone copolyol (10% in cyclomethicone) (40.52 g); C12-15 alkyl benzoate (60.17 g); and cyclomethicone (49.83g) and mixing them at 500 rpm until the mixture was homogeneous to form Phase A. Phase B was made by combining an antiperspirant active (Westchlor ZR 35B 30% PG solution) (152.07 g), Polysorbate 80 (1.30 g), propylene glycol (146.82 g), ethanol (95% alc.) (45.06 g), and fragrance (5.02 g). Phase B was added to Phase A with stirring and the composition was allowed to sit overnight and placed in a package with a porous applicator.

REFERENCE COUNT: 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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ANSWER 10 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                        1999:722376 CAPLUS
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DOCUMENT NUMBER: 132:298673

TITLE: Nebulizer-compatible liquid formulations for pulmonary delivery of glucocorticoids: pre-formulation studies Klyashchitsky, B.; Saidi, Z.; Saar, A.; Sedlak, D.; AUTHOR(S):

Szymkowiak, J.; Owen, A.

CORPORATE SOURCE: LDS Technologies, Inc., Boothwyn, PA, 19061, USA SOURCE: Proceedings of the International Symposium on Controlled Release of Bioactive Materials (

1999), 26th, 565-566

CODEN: PCRMEY; ISSN: 1022-0178 PUBLISHER: Controlled Release Society, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

The determination of solubility in various solvents, oils, and surfactants and AB stability evaluation was valuable in the pre-formulation of glucocorticoid liquid composition development.

L5 ANSWER 11 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1998:661508 CAPLUS DOCUMENT NUMBER: 129:281013

TITLE: Pharmaceutical compositions containing lamivudine and

a preservative

INVENTOR(S): INVENTOR(S): Nguyen, Ngoc-Anh Thi; Casey, Warren M.
PATENT ASSIGNEE(S): Glaxo Group Limited, UK
SOURCE: PCT Int. Appl., 15 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. Di MO 9842321 A2 19981001 MO 1998-EP1626 15 WO 9842321 A3 19990107 W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP,	9980320 < CZ, DE, KE, KG, MW, MX,
W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU,	CZ, DE, KE, KG, MW, MX,
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DK. EE. ES. FI. GB. GE. GH. GM. GW. HU. ID. IL. IS. JP.	MW, MX,
KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN,	
NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM,	TR, TT,
UA, UG, US, UZ, VN, YU, ZW	
RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK,	
FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, GA, GN, ML, MR, NE, SN, TD, TG	C1, CM,
GA, GR, ML, MR, NE, SN, TD, TG TM 536403 B 20030611 TW 1998-87103841 19 ZA 9902367 A 19990920 ZA 1998-2367 11 CA 2286126 A1 19981001 CA 1998-2286126 19 CA 2286126 C 20030812 AU 9972084 A 19981020 AU 1998-72084 19 AU 728461 B2 20010111 US 6004968 A 19991221 US 1998-44896 19 EP 969815 A2 20000112 EP 1998-919120 19 EP 969815 B1 20050511 EP 371 BE CH DE DK ES FR GB, GR, TT, LI, LU, NL, SE,	0000016
77 0002267 7 10000020 77 1000 2267 10	9980316 <
Ch 2206126 h 19990920 hh 1990-2307 1:	2200313 <
CA 2200120 A1 19901001 CA 1990-2200120 1:	9900320 <
ATT 9872084 A 19981020 ATT 1998-72084 19	9980320 <
AU 728461 B2 20010111	3300320 1
US 6004968 A 19991221 US 1998-44896 19	9980320 <
EP 969815 A2 20000112 EP 1998-919120 19	9980320 <
EP 969815 B1 20050511	
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE,	MC, PT,
IE, SI, LT, LV, FI, RO	
IE, SI, LT, LV, FI, RO BR 9808060 A 20000308 BR 1998-8060 19 EE 9900440 A 20000417 EE 1999-440 19 EE 3996 B1 20030415 IL 131917 A 20010111 IL 1998-131917 19 HU 2000002982 A2 20010129 HU 2000-2982 19 HU 2000002982 A3 20011228 HU 225600 B1 20070502 JP 2001501974 T 20010213 JP 1998-544425 19 JP 3264937 B2 20020311 JP 3264937 B2 20020311 NZ 337798 A 20010330 NZ 1998-337798 19 IL 138098 A 20030112 IL 1998-138098 19	9980320 <
EE 9900440 A 20000417 EE 1999-440 19	9980320 <
EE 3996 B1 20030415	
IL 131917 A 20010111 IL 1998-131917 19	9980320 <
HU 2000002982 A2 20010129 HU 2000-2982 19	9980320 <
HU 2000002982 A3 20011228	
HU 225600 B1 20070502	
JP 2001501974 T 20010213 JP 1998-544425 19	9980320 <
JP 3264937 B2 20020311	
NZ 337/98 A 20010330 NZ 1998-337/98 15	9980320 <
AP 1141 A 20030129 AP 1999-1657 19	9980320 <
	9900320 <
CV 202/17 DC 20020701 CV 1000_1200 10	0000320 <
W: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW SK 283417 B6 20030701 SK 1999-1299 19 AT 295150 T 20050515 AT 1998-919120 19 PT 969815 T 20050729 PT 1998-919120 19	9980320 \
PT 969815 T 20050313 RT 1550 515120 1:	9980320
ES 2239802 T3 20051001 ES 1998-919120 19	9980320
PI. 190505 B1 20051230 PI. 1998-336038 19	9980320
CZ 298008 B6 20070523 CZ 1999-3403 19	9980320
HR 980154 B1 20020630 HR 1998-154 19	9980323 <
IN 1998CA00479 A 20050318 IN 1998-CA479 19	9980323
SK 283417 B6 20030701 SK 1999-1299 1: AT 295150 T 20050515 AT 1998-919120 1: PT 969815 T 20050729 PT 1998-919120 1: ES 2239802 T3 20051001 ES 1998-919120 1: PL 190505 B1 20051230 PL 1998-336038 1: CZ 298008 B6 20070523 CZ 1999-3403 1: EN 980154 B1 20020630 HR 1998-1544 1: IN 1998CA00479 A 20050318 IN 1998-CA479 1: MX 9908690 A 20000131 MX 1999-6690 1:	9990922 <

NO 9904619 BG 64690 HK 1022853 PRIORITY APPLN. INFO.	A B1 A1	19991123 20051230 20050909	NO 1999-4619 BG 1999-103818 HK 2000-102154 US 1997-42353P GB 1997-6295 IL 1998-131917	A3	19990923 < 19991018 20000407 19970324 19970326 19980320
			WO 1998-EP1626	W	19980320

AB Oral antiviral formulations containing lamivudine, substantially free of EtOH and EDTA, contain parabens at pH >5.5 as preservatives. Thus, a liquid formulation contained lamivudine 10.00, sucrose 200.0, Me paraben 1.50 kg, Pr paraben 180, artificial strawberry flavor 800, artificial banana flavor 600, Na citrate-2H2O 11, anhydrous citric acid

1 g, NaOH or HCl to pH 6.0, propylene glycol 19.4, and H2O to 1000 L. This composition remained free from growth of inoculated bacteria, yeasts, and molds for 14-28 days.

L5 ANSWER 12 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1998:635637 CAPLUS

DOCUMENT NUMBER: 129:265476

TITLE: Stable particle in liquid formulations comprising

sugar glass

INVENTOR(S): Roser, Bruce Joseph; Sen, Shevanti Devika

PATENT ASSIGNEE(S): Eastbridge Ltd., UK SOURCE: PCT Int. Appl., 40 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

										APPLICATION NO.									
WO		188			A2		1998	0924									318 <		
WO															011				
	₩:										BY,								
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											LV,								
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							YU,												
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CA	22936	682			A1		1998	0924		CA 1	998-	2293	682		1	9980	318 <		
CA	22936	682			С		2007	0116											
AU	9865	101			A.		1998	1012		AU 1	998-	6510	1		1	9980	318 <		
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EP	10070																		
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BR	98089	920			A												318 <		
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PRIORIT:	Y APPI	LN.	INFO	.:							997-								
										WO 1	998-	GB81	7		W 1	9980	318		

AB A stable particle in liquid formulation comprising a discontinuous phase of microparticles is suspended in a continuous phase which is a non-aqueous liquid,

preferably biocompatible in which the microparticles are insol. The microparticles comprise finely powdered sugar glass selected from the group consisting of trehalose, palatinit, glucopyranosyl sorbitol, glucopyranosyl mannitol, lactitol and monosaccharide alcs. such as mannitol and inositol, holding at least one biomol, product, the biomol. product in the sugar glass either being in stable solid solution or being itself in suspension in the sugar glass. A monodisperse single-particle suspension of microparticles can be produced in the non-aqueous continuous liquid phase by inclusion in the continuous phase of at least one surfactant having a low or very low HLB. A solution containing trehalose 0.6, sodium sulfate 0.35 M, bovine serum albumin 0.75, zinc chloride 1, magnesium chloride 1 mM, and alkaline phosphatase 40 units/mM was spray dried. When the powder was stored at 37°, there was no loss of enzyme activity over 84 days of storage.

L5 ANSWER 13 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1997:650275 CAPLUS

DOCUMENT NUMBER: 127:298754

TITLE: Compositions comprising an HIV protease inhibitor such as VX 478 and a water soluble vitamin e compound such

as vitamin E-TPGS

INVENTOR(S): Roy, Arup K.; Tillman, Lloyd Gary
PATENT ASSIGNEE(S): Glaxo Group Limited, UK; Roy, Arup K.; Tillman, Lloyd

Gary SOURCE: PCT Int. Appl., 22 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PAT	TENT I	NO.			KIN	D	DATE			APPLICATION NO.						DATE			
															19970321 <				
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	RW:																		
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TD	2000 3117 3316	726	04		D 2		2000	1210		JP	1997	-3340	1/		1	9910	221	·	
N7	3316	15			7		2000	1210		MZ	1997.	-3316	45		1	9970	321		
Thi	4554	91			D		2000	0220		T167	1997	-0610	3607		1		321		
C7	2899	58			B6		2001	0515		C7	1998	-3035	5007		1		321		
ΔT	4554 2899 2306	02			т		2002	0115		ΔT	1997	-9142	87		1		321		
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20030314
    AP 1150
                      A
                                      AP 1998-1343 19970321 <--
       W: GH, GM, KE, LS, MW, SD, SZ, UG, ZW
    IL 126185 A 20030529 IL 1997-126185
                                                           19970321 <--
    ES 2190528
                      Т3
                           20030801 ES 1997-914287
                                                            19970321
    EE 4093
                      B1 20030815 EE 1998-323
                                                            19970321
    PL 187919
                      B1 20041130 PL 1997-328916
                                                            19970321
                     B6 20041201 SK 1998-1269
A 20050311 IN 1997-DE727
    SK 284244
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    IN 1997DE00727
                                                            19970321
                     B1 20050630 RO 1998-1407
A 19981120 NO 1998-4386
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    NO 317639
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                                                            19980921 <--
    BG 64457
                      B1 20050331 BG 1998-102838
A1 20030711 HK 1999-102135
                                                            19981012
    HK 1016896
                                                            19990512
PRIORITY APPLN. INFO.:
                                       US 1996-13893P
                                                        P 19960322
                                       GB 1996-6372
                                                        A 19960326
                                       US 1997-820848
                                                        A 19970320
                                       WO 1997-EP1438
                                                        W 19970321
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AB Pharmaceutical formulations containing HIV protease inhibitors, specifically including 3S-[3R*(1R*,2S*)]-[3-[((4-aminophenyl)sulfonyl)(2-methylpropyl)-amino]-2-hydroxy-1-(phenylmethyl)propyl]carbamic acid, tetrahydro-3-furanyl ester (alternatively known as VX 478 or 141W94) (I), and a tocopherol, and their use in medical therapy are described. A liquid formulation was prepared containing I 150.0, α-tocopheryl PEG succinate (TPGS) 400.0, PEG 400 200.5, and propylene glycol 39.5 mg/capsule.

L5 ANSWER 14 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1996:313848 CAPLUS

DOCUMENT NUMBER: 124:352705

TITLE: Tastemasked liquid pharmaceuticals containing sugars and hydrogenated maltose and polyhydroxy alcohols

INVENTOR(S): Lienhop, Keith S.; Cuca, Robert C.; Riley, Thomas Charles, Jr.; Levinson, R. Saul

PATENT ASSIGNEE(S): Kv Pharmaceutical Corporation, USA

SOURCE: PCT Int. Appl., 24 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

LANGUAGE: Engl FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

by

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9603976	A1 199602	15 WO 1995-US9709	19950801 <
W: AU, CA, JP,	MX, NZ		
RW: AT, BE, CH,	DE, DK, ES, F	R, GB, GR, IE, IT, LU, MC	, NL, PT, SE
CA 2172807	A1 199602	15 CA 1995-2172807	19950801 <
CA 2172807	C 199910	12	
AU 9531548	A 199603	04 AU 1995-31548	19950801 <
US 5730997	A 199803	24 US 1996-712436	19960911 <
PRIORITY APPLN. INFO.:		US 1994-282495	A 19940801
		WO 1995-US9709	W 19950801

AB A substantially tasteless liquid pharmaceutical delivery system containing an active material and a high osmolarity aqueous system comprising (1) water; (2) about 20% to about 45% by weight sugar derivative; (3) about 10% to about 15%

weight hydrogenated maltose syrup; and (4) about 0% to about 35% by weight polyhydroxy alc. A tastemasked liquid formulation contained diphenhydramine.HCl 0.2111, water 16.4624, sorbitol solution

contained diphenhydramine.HCl 0.2111, water 16.4624, sorbitol solutio 41.8179, maltitol solution 13.9287, propylene glycol

25.8670, sodium gluconate 0.1857, citric acid 0.2111, saccharin sodium

0.1013, magnasweet-180 0.0422, Me paraben 0.844, Pr paraben 0.0152, colors 0.0177, and flavor 1.0553%.

L5 ANSWER 15 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1994:452376 CAPLUS

DOCUMENT NUMBER: 121:52376

TITLE: Stabilized isothiazolone liquid formulation INVENTOR(S): Sano, Yoichi; Tsuji, Katsuji; Katayama, Sakae

PATENT ASSIGNEE(S): Katavama Chemical Inc., Japan

SOURCE: U.S., 5 pp. Cont. of U.S. Ser. No. 745,250, abandoned.

CODEN: USXXAM DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. ---------19940426 US 1992-970231 19921030 US 1991-745250 B1 19910814 19921030 <--US 5306725 PRIORITY APPLN. INFO.: MARPAT 121:52376

OTHER SOURCE(S):

A Stabilized isothiazolone liquid formulation including: an isothiazolone AB compound represented by the formula (I): (where X represents a H atom or halogen atom, and Y represents a lower alkyl group), and a mixed solvent containing 50-99.9 weight% of a glycol type solvent and 50-0.1 weight% of an amide-type compound represented by the formula R1CONR2R3, where R1 represents a H atom or a lower alkyl group, R2 and R3 each represent a lower alkyl group, R1 and R3 each represent a lower alkyl group, R1 may bond to R2 or R3 to form a nitrogen-containing heterocycle, the compound of the formula I being dissolved in the mixed solvent of which amount is at least sufficient to dissolve it.

L5 ANSWER 16 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1993:598571 CAPLUS

DOCUMENT NUMBER: 119:198571

TITLE: Stabilized aqueous liquid formulations of phytase and their use in feed preparation for monogastric animals INVENTOR(S): Barendse, Rudolfus Carolus Mari: Van Doesum, Johannes

Henricus; Gouwens, Jacob; Van Paridon, Petrus Andreas

PATENT ASSIGNEE(S): Gist-Brocades N.V., Neth. SOURCE: PCT Int. Appl., 23 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9316175	A1	19930819	WO 1993-EP356	19930212 <
W. AH BB BG	BR CA	CZ ET HII	JP KP KR LK MG N	IN MW NO

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        RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
            BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, SN, TD, TG
    AII 9336284
                               19930903 AU 1993-36284
                         A
                                                                  19930212 <--
    EP 626010
                         A1
                               19941130
                                          EP 1993-905244
                                                                 19930212 <--
        R: DE, DK, NL
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                                                                  19940810 <--
PRIORITY APPLN. INFO.:
                                           EP 1992-200414
                                                              A 19920213
                                           WO 1993-EP356
                                                              A 19930212
    A stabilized liquid formulation of phytase contains a stabilizing agent,
    i.e. urea 1-10 weight/weight% or water-soluble polyol, such as sorbitol or
    glycerol. A feed composition for monogastric animals is prepared by treating
the
    feed with the stabilized phytase formulation. The treatment releases P
    from the phytate, making it available to the animal.
   ANSWER 17 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                       1965:50780 CAPLUS
DOCUMENT NUMBER:
                        62:50780
ORIGINAL REFERENCE NO.: 62:8946c-d
TITLE:
                        Metal-acid complexes with members of the tetracycline
                        family. II. Development of stable preconstituted
                        parenteral formulations
                        Remmers, Edward G.; Barringer, William C.; Sieger,
AUTHOR(S):
                        George M.; Doerschuk, Albert P.
CORPORATE SOURCE:
                        Am. Cyanamid Co., Pearl River, NY
SOURCE:
                        Journal of Pharmaceutical Sciences (1964),
                        53(12), 1534-6
                        CODEN: JPMSAE; ISSN: 0022-3549
DOCUMENT TYPE:
                        Journal
LANGUAGE:
                        English
AB cf. CA 62, 3888h. Al-Ca-gluconate complexes of tetracycline and
    6-demethylchlortetracycline were prepared by previously described methods
    (loc. cit.) and made into stable liquid formulations
    suitable for intramuscular and intravenous use by solution in aqueous
    propylene glycol (I). The formulations were well
    tolerated at therapeutic levels and gave adequate blood levels. Prepns.
    containing the 1:3:1:6 (molar ratio) antibiotic-Al-Ca-gluconate complex in
    50-75% I at pH 8.5 were the most satisfactory and retained initial
    potencies at both room and elevated temps. for prolonged periods.
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